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Amendments to the Claims

The following listing of claims will replace all prior versions, and listings, of claims in

the present application:

1. (Currently Amended) A solid corrosion-inhibiting conversion coating formed on a

substrate metal, the conversion coating comprising a rare earth element and an  $\underline{\text{inorganic}}$  valence

stabilizer combined to form a rare earth/valence stabilizer complex within the solid corrosion-

inhibiting conversion coating, wherein the rare earth element is selected from cerium, praseodymium, terbium, or a combination thereof, and at least one rare earth element is in the

tetravalent oxidation state in the rare earth/valence stabilizer complex in the solid corrosion-

inhibiting conversion coating, and wherein the valence stabilizer consists essentially of an

inorganic valence stabilizer.

2. (Original) The conversion coating of claim 1 wherein the rare earth/valence stabilizer

complex has a solubility in water of between about 5 x  $10^{-1}$  and about 1 x  $10^{-5}$  moles per liter of

cerium, praseodymium, or terbium at about 25°C and about 760 Torr.

 $3. \ (Original) \quad The \ conversion \ coating \ of \ claim \ 2 \ wherein \ the \ solubility \ of \ the \ rare \ earth/valence$ 

stabilizer complex in water is between about 5 x  $10^{-2}$  and about 5 x  $10^{-5}$  moles per liter of cerium,

praseodymium, or terbium at about 25°C and about 760 Torr.

4. (Original) The conversion coating of claim 1 wherein there is an electrostatic barrier layer

around the rare earth/valence stabilizer complex in aqueous solution.

5. (Original) The conversion coating of claim 1 wherein the rare earth/valence stabilizer

complex acts as an ion exchange agent towards corrosive ions.

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- 6. (Original) The conversion coating of claim 1 wherein the conversion coating is between about 25 and about 10,000 nanometers thick.
- 7. (Original) The conversion coating of claim 6 wherein the conversion coating is between about 100 and about 500 nanometers thick.
- 8. (Original) The conversion coating of claim 1 wherein the conversion coating has a morphology which enhances adhesion of a coating applied over the conversion coating.
- 9. (Original) The conversion coating of claim 1 wherein the rare earth/valence stabilizer complex has a central cavity containing a cerium, praseodymium, or terbium ion and an additional ion.
- 10. (Original) The conversion coating of claim 9 wherein the additional ion is B<sup>+3</sup>, Al<sup>+3</sup>, Si<sup>+4</sup>, P<sup>+5</sup>, Ti<sup>+4</sup>, V<sup>+5</sup>, Vi<sup>+4</sup>, Cr<sup>+6</sup>, Cr<sup>+3</sup>, Mn<sup>+4</sup>, Mn<sup>+3</sup>, Mn<sup>+2</sup>, Fe<sup>+3</sup>, Fe<sup>+2</sup>, Co<sup>+2</sup>, Co<sup>+3</sup>, Ni<sup>+2</sup>, Ni<sup>+3</sup>, Ni<sup>+4</sup>, Cu<sup>+2</sup>, Cu<sup>+3</sup>, Zn<sup>+2</sup>, Ga<sup>+3</sup>, Ge<sup>+4</sup>, As<sup>+5</sup>, As<sup>+5</sup>, or Zr<sup>+4</sup>
- 11. (Canceled)
- 12. (Previously Presented) The conversion coating of claim 1 wherein the inorganic valence stabilizer is selected from molybdates, tungstates, vanadates, niobates, tantalates, tellurates, periodates, iodates, carbonates, antimonates, stannates, phosphates, nitrates, bromates, sulfates, titanates, zirconates, bismuthates, germanates, arsenates, selenates, borates, aluminates, silicates, or combinations thereof.

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13. (Original) The conversion coating of claim 12 wherein the valence stabilizer is the inorganic valence stabilizer selected from molybdates, tungstates, vanadates, niobates, tantalates, tellurates, periodates, iodates, carbonates, antimonates, stannates, phosphates, nitrates, bromates, sulfates, or combinations thereof.

14-36. (Canceled)

- 37. (Previously Presented) The conversion coating of claim 1 wherein the rare earth/valence stabilizer complex further comprises a solubility control agent.
- 38. (Original) The conversion coating of claim 37 wherein the solubility control agent is a cationic solubility control agent or an anionic solubility control agent.
- 39. (Original) The conversion coating of claim 38 wherein the solubility control agent is the cationic solubility control agent selected from H\*; Li\*; Na\*; K\*; Rb\*; Cs\*; NH4\*; Mg\*2; Ca\*2; Sr\*2; Be\*2; Ba\*2; Y\*3; La\*3; Ce\*3; Ce\*4; Nd\*3; Pr\*3; Se\*3; Sm\*3; Eu\*3; Eu\*2; Gd\*3; Tb\*3; Dy\*3; Ho\*3; Ev\*3; Tm\*3; Yb\*3; Lu\*3; Ti\*4; Zr\*4, Ti\*3; Hf\*4; Nb\*5; Ta\*5; Nb\*4; Ta\*4, V\*5; V\*4, V\*3; Mo\*6; W\*6; Mo\*5; W\*5; Mo\*4; W\*4; Cr\*3; Mn\*3; Mn\*3; Mn\*4; Fe\*2; Fe\*3; Co\*2; Co\*3; Ni\*2; Ni\*3; Ni\*4; Ru\*2; Ru\*3; Ru\*4; Rh\*3; Ir\*3; Rh\*2; Ir\*2; Pa\*4; Pt\*4; Pd\*2; Pt\*2; Os\*4; Cu\*2; Cu\*3; Ag\*; Ag\*2; Ag\*3; Au\*2; Au\*2; Au\*3; Zn\*2; Cd\*2; Hg\*1; Hg\*2; Al\*3; Ga\*3; Ga\*; In\*3; In\*; Tl\*3; Tl\*; Ge\*4; Ge\*2; Sn\*4; Sn\*2; Pb\*4; Pb\*2; Sb\*3; Sb\*5; As\*3; As\*5; Bi\*3; Bi\*5; organic compounds containing at least one N\* site; organic compounds containing at least one sulfonium site; organic compounds containing at least one solutionium site; organic compounds co

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formula  $NR_4^+$ , where R is an alkyl, aromatic, or acyclic organic constituent; or combinations thereof.

40. (Original) The conversion coating of claim 39 wherein the cationic solubility control agent is selected from H<sup>+</sup>; Li<sup>+</sup>; Na<sup>+</sup>; K<sup>+</sup>; Rb<sup>+</sup>; Cs<sup>+</sup>; NH<sub>4</sub><sup>+</sup>; Mg<sup>+2</sup>; Ca<sup>+2</sup>; Sr<sup>+2</sup>; Y<sup>+3</sup>; La<sup>+3</sup>; Ce<sup>+3</sup>; Ce<sup>+4</sup>; Nd<sup>+3</sup>; Pr<sup>+3</sup>; Sc<sup>+3</sup>; Sm<sup>+3</sup>; Eu<sup>+3</sup>; Eu<sup>+2</sup>; Gd<sup>+2</sup>; Tb<sup>+3</sup>; Dy<sup>+3</sup>; Ho<sup>+3</sup>; Er<sup>+3</sup>; Tm<sup>+2</sup>; Yb<sup>+3</sup>; Lu<sup>+3</sup>; Ti<sup>+4</sup>; Zr<sup>+4</sup>; Ti<sup>+3</sup>; Hf<sup>+4</sup>; Nb<sup>+5</sup>; Ta<sup>+5</sup>; Nb<sup>+4</sup>; Ta<sup>+4</sup>; Mo<sup>+6</sup>; W<sup>+6</sup>; Mo<sup>+5</sup>; Wh<sup>-5</sup>; Mo<sup>+4</sup>; Wh<sup>+4</sup>; Mn<sup>+2</sup>; Mn<sup>+3</sup>; Mn<sup>+4</sup>; Fe<sup>+2</sup>; Fe<sup>+3</sup>; Co<sup>+2</sup>; Co<sup>+3</sup>; Ru<sup>+2</sup>; Ru<sup>+3</sup>; Ru<sup>+4</sup>; Rh<sup>+3</sup>; Ir<sup>+3</sup>; Rh<sup>+2</sup>; Ir<sup>2</sup>; Pd<sup>+4</sup>; Pt<sup>+4</sup>; Pd<sup>+2</sup>; Pt<sup>+2</sup>; Cu<sup>+</sup>; Cu<sup>+2</sup>; Cu<sup>+3</sup>; Ag<sup>+</sup>; Ag<sup>+2</sup>; Ag<sup>+3</sup>; Au<sup>+</sup>; Au<sup>+3</sup>; Zn<sup>+2</sup>; Al<sup>+3</sup>; Ga<sup>+3</sup>; Ga<sup>+3</sup>; Ga<sup>+3</sup>; Ga<sup>+4</sup>; Pf<sup>+2</sup>; Pf<sup>+2</sup>; Sh<sup>+4</sup>; Sn<sup>+2</sup>; Sh<sup>+3</sup>; Sh<sup>+5</sup>; Bi<sup>+3</sup>; Bi<sup>+5</sup>; organic compounds containing at least one N<sup>+</sup> site; organic compounds containing at least one sulfonium site; organic compounds containing at least one sulfonium site; organic compounds having a formula NR<sub>4</sub><sup>+</sup>, where R is an alkyl, aromatic, or acyclic organic constituent; or combinations thereof.

41. (Withdrawn) The conversion coating of claim 38 wherein the solubility control agent is the anionic solubility control agent selected from fluorotitanates, chlorotitanates, fluorozirconates, chlorozirconates, fluoroniobates, chloroniobates, fluorotantalates, chlorotantalates, molybdates, tungstates, permanganates, fluoromanganates, chloromanganates, fluoroferrates, chloroferrates, fluorocobaltates, chlorocobaltates, fluorozincates, chlorozincates, borates, fluorostannates, fluoroaluminates, carbonates, silicates, fluorosilicates, fluorostannates, nitrates, nitrites, azides, cyanamides, phosphates, phosphinites, fluorophosphates, thiophosphates, thiophosphates, thiophosphates, thiophosphates, thiophosphates, thiophosphates, thiophosphates, thiophosphates, fluoroantimonates, chloroantimonates, sulfates, sulfites, sulfonates, thiosulfates, dithionites, dithionates, fluorosulfates, tellurates, fluorides, chlorates, perchlorates, bromides, bromates, iodides, iodates, periodates, heteropolyanions, ferricyanides, ferrocyanides,

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cyanocobaltates, cyanocuprates, cyanomanganates, cyanatos ferrates, cyanatocobaltates, cyanatocuprates, cyanatomanganates, thiocyanates, thiocyanatoferrates, thiocyanatocobaltates, thiocyanatocuprates, thiocyanatomanganates, cyanamides, cyanamidoferrates, cyanamidocobaltates, cyanamidocuprates, cyanamidomanganates, nitritoferrates. nitritocobaltates, azides, (thio)carboxylates, di(thio)carboxylates, tri(thio)carboxylates, tetra(thio)carboxylates, (thio)phenolates, di(thio)phenolates, tri(thio)phenolates, tetra(thio)phenolates, (thio)phosphonates, di(thio)phosphonates, tri(thio)phosphonates, (thio)phosphonamides, di(thio)phosphonamides, tri(thio)phosphonamides, amino(thio)phosphonates, diamino(thio)phosphonates, triamino(thio)phosphonates, imino(thio)phosphonates, diimino(thio)phosphonates, (thio)sulfonates, di(thio)sulfonates, tri(thio)sulfonates, (thio)sulfonamides, di(thio)sulfonamides, tri(thio)sulfonamides, amino(thio)sulfonates, diamino(thio)sulfonates, triamino(thio)sulfonates, imino(thio)sulfonates, diimino(thio)sulfonates, (thio)borates, di(thio)borates, (thio)boronates, organic silicates, stibonates, cyanides, cyanochromates, cyanonickelates, cyanatochromates, cyanatonickelates, thiocvanatochromates, thiocvanatonickelates, cyanamidochromates, cyanamidonickelates, nitritonickelates, arsonates, diarsonates, triarsonates, organic selenates, diselenates, triselenates, arsenates, arsenites, fluoroarsenates, chloroarsenates, selenates, selenites, fluorothallates, chlorothallates, iodomercury anions, chloromercurates, bromomercurates, osmates, fluoronickelates, chromates, Reinecke's salt, vanadates, or combinations thereof.

42. (Withdrawn) The conversion coating of claim 41 wherein the anionic solubility control agent is selected from fluorotitanates, chlorotitanates, fluorozirconates, chlorozirconates, fluoroniobates, chloroniobates, fluorotantalates, chlorotantalates, molybdates, tungstates, permanganates, fluoromanganates, chloromanganates, fluoroferrates, chloroferrates, fluorocobaltates, chlorocobaltates, fluorozincates, chlorozincates, borates, fluorostannates, fluoroaluminates, chloroaluminates, carbonates, fluorosilicates, fluorosilicates, fluorostannates, nitrates, nitrites, azides, cyanamides, phosphates, phosphinites, phosphonates, phosphinites,

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thiophosphates, thiophosphites, thiophosphonates, thiophosphinites, fluorophosphates, fluoroantimonates, chloroantimonates, sulfates, sulfites, sulfonates, thiosulfates, dithionites, dithionates, fluorosulfates, tellurates, fluorides, chlorides, chlorates, perchlorates, bromides, bromates, iodides, iodates, periodates, heteropolyanions, ferricyanides, ferrocyanides, cyanocobaltates, cyanocuprates, cyanomanganates, cyanatoferrates, cyanatocobaltates, cyanatocuprates, cyanatomanganates, thiocyanates, thiocyanatoferrates, thiocyanatocobaltates, thiocyanatocuprates, thiocyanatomanganates, cyanamides, cyanamidoferrates, cyanamidocobaltates, cyanamidocuprates, cyanamidomanganates, nitritoferrates, nitritocobaltates, azides, (thio)carboxylates, di(thio)carboxylates, tri(thio)carboxylates, tetra(thio)carboxylates, (thio)phenolates, di(thio)phenolates, tri(thio)phenolates, tetra(thio)phenolates, (thio)phosphonates, di(thio)phosphonates, tri(thio)phosphonates, (thio)phosphonamides, di(thio)phosphonamides, tri(thio)phosphonamides, amino(thio)phosphonates, diamino(thio)phosphonates, triamino(thio)phosphonates, imino(thio)phosphonates, diimino(thio)phosphonates, (thio)sulfonates, di(thio)sulfonates, tri(thio)sulfonates, (thio)sulfonamides, di(thio)sulfonamides, tri(thio)sulfonamides. amino(thio)sulfonates, diamino(thio)sulfonates, triamino(thio)sulfonates, imino(thio)sulfonates, diimino(thio)sulfonates, (thio)borates, di(thio)borates, (thio)boronates, organic silicates, stibonates, or combinations thereof.

- 43. (Original) The conversion coating of claim 1 wherein the conversion coating is colored.
- 44. (Original) The conversion coating of claim 43 further comprising an agent which improves color-fastness of the conversion coating.
- 45. (Original) The conversion coating of claim 44 wherein the agent which improves color-fastness is selected from an active UV blocker, a passive UV blocker, a brightener, or a combination thereof.

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46. (Original) The conversion coating of claim 45 wherein the agent which improves color-

fastness is the active UV blocker selected from carbon black, graphite, phthalocyanines, or

combinations thereof.

47. (Original) The conversion coating of claim 45 wherein the agent which improves color-

fastness is the passive UV blocker selected from titanium oxide, tin oxide, lead oxide, silicon

oxide, silicates, aluminosilicates, or combinations thereof.

48. (Original) The conversion coating of claim 45 wherein the agent which improves color-

fastness is the brightener selected from sulfonic acids, sulfonates, sulfonamides, sulfinic acids,

sulfinates, sulfones, cyanides, nonionic surfactants, or combinations thereof.

49. (Original) The conversion coating of claim 43 wherein the color is formed by a dye selected from vat dyes, mordant dyes, lake dyes, disperse dyes, azo dyes, triazene dyes, triphenylmethane

dyes, azine dyes, formazan dyes, phthalocyanine dyes, Schiff Base dyes, naturally-occurring

dyes, inorganic pigments, or combinations thereof.

50-162 (Canceled)

163. (Currently Amended) A solid corrosion-inhibiting conversion coating formed on a

substrate metal, the conversion coating comprising a rare earth element and an inorganic valence

stabilizer combined to form a rare earth/valence stabilizer complex within the solid corrosion-

inhibiting conversion coating, wherein

the rare earth element is selected from cerium, praseodymium, terbium, or a combination

thereof,

at least one rare earth element is in the tetravalent oxidation state in the rare earth/valence

stabilizer complex in the solid corrosion-inhibiting conversion coating, and

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the rare earth/valence stabilizer complex is sparingly soluble in water at about 25°C and about 760 Torr, and

wherein the valence stabilizer consists essentially of an inorganic valence stabilizer.

164. (Canceled)

165. (Previously Presented) A solid corrosion-inhibiting conversion coating formed on a substrate metal, the conversion coating comprising a rare earth element and a valence stabilizer combined to form a rare earth/valence stabilizer complex within the solid corrosion-inhibiting conversion coating, wherein the rare earth element is selected from cerium, praseodymium, terbium, or a combination thereof, and at least one rare earth element is in the tetravalent oxidation state, wherein the rare earth/valence stabilizer complex has a central cavity containing a cerium, praseodymium, or terbium ion and an additional ion, wherein the additional ion is B<sup>13</sup>, Al<sup>13</sup>, Si<sup>14</sup>, P<sup>15</sup>, Ti<sup>14</sup>, V<sup>15</sup>, V<sup>14</sup>, Cr<sup>16</sup>, Cr<sup>23</sup>, Mn<sup>14</sup>, Mn<sup>23</sup>, Mn<sup>22</sup>, Fe<sup>23</sup>, Fe<sup>23</sup>, Co<sup>23</sup>, Co<sup>33</sup>, Ni<sup>23</sup>, Ni<sup>23</sup>, Ni<sup>24</sup>, Cu<sup>23</sup>, Cu<sup>23</sup>, Zn<sup>23</sup>, Ga<sup>23</sup>, Ge<sup>24</sup>, As<sup>25</sup>, As<sup>25</sup>, or Zr<sup>24</sup>.

166. (Previously Presented) The conversion coating of claim 165 wherein the rare earth/valence stabilizer complex has a solubility in water of between about 5 x 10<sup>-1</sup> and about 1 x 10<sup>-5</sup> moles per liter of cerium, praseodymium, or terbium at about 25°C and about 760 Torr.

167. (Previously Presented) The conversion coating of claim 166 wherein the solubility of the rare earth/valence stabilizer complex in water is between about  $5 \times 10^{-2}$  and about  $5 \times 10^{-5}$  moles per liter of cerium, praseodymium, or terbium at about 25°C and about 760 Torr.

168. (Previously Presented) The conversion coating of claim 165 wherein there is an electrostatic barrier layer around the rare earth/valence stabilizer complex in aqueous solution.

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 $169. \ (Previously\ Presented) \quad The\ conversion\ coating\ of\ claim\ 165\ wherein\ the\ rare$ 

earth/valence stabilizer complex acts as an ion exchange agent towards corrosive ions.

170. (Previously Presented) The conversion coating of claim 165 wherein the conversion

coating is between about 25 and about 10,000 nanometers thick.

171. (Previously Presented) The conversion coating of claim 170 wherein the conversion

coating is between about 100 and about 500 nanometers thick.

172. (Previously Presented) The conversion coating of claim 165 wherein the conversion

coating has a morphology which enhances adhesion of a coating applied over the conversion

coating.

173. (Previously Presented) The conversion coating of claim 165 wherein the valence stabilizer

is an inorganic valence stabilizer selected from molybdates, tungstates, vanadates, niobates,

tantalates, tellurates, periodates, iodates, carbonates, antimonates, stannates, phosphates, nitrates,

 $bromates,\, sulfates,\, titanates,\, zirconates,\, bismuthates,\, germanates,\, arsenates,\, selenates,\, borates,\, titanates,\, tita$ 

aluminates, silicates, or combinations thereof.

174. (Previously Presented) The conversion coating of claim 173 wherein the valence stabilizer

is the inorganic valence stabilizer selected from molybdates, tungstates, vanadates, niobates,

tantalates, tellurates, periodates, iodates, carbonates, antimonates, stannates, phosphates, nitrates,

bromates, sulfates, or combinations thereof.

175. (Previously Presented) The conversion coating of claim 165 wherein the rare earth/valence

stabilizer complex further comprises a solubility control agent.

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176. (Previously Presented) The conversion coating of claim 175 wherein the solubility control agent is a cationic solubility control agent or an anionic solubility control agent.

177. (Previously Presented) The conversion coating of claim 176 wherein the solubility control agent is the cationic solubility control agent selected from H<sup>+</sup>; Li<sup>+</sup>; Na<sup>+</sup>; K<sup>+</sup>; Rb<sup>+</sup>; Cs<sup>+</sup>; NH<sub>4</sub><sup>+</sup>; Mg<sup>+</sup><sup>2</sup>; Ca<sup>+</sup><sup>2</sup>; Sr<sup>+</sup><sup>2</sup>; Ba<sup>+</sup><sup>2</sup>; Y<sup>+</sup><sup>3</sup>; La<sup>+</sup><sup>3</sup>; Ca<sup>+</sup><sup>3</sup>; Ce<sup>+</sup><sup>4</sup>, Nd<sup>+</sup><sup>3</sup>; Pr<sup>+</sup><sup>3</sup>; Sc<sup>+</sup><sup>3</sup>; Sm<sup>+</sup><sup>3</sup>; Eu<sup>+</sup><sup>3</sup>; Eu<sup>+</sup><sup>2</sup>; Gd<sup>+</sup><sup>3</sup>; Tb<sup>+</sup><sup>3</sup>; Dy<sup>+</sup><sup>3</sup>; Ho<sup>+</sup><sup>3</sup>; Tm<sup>+</sup><sup>3</sup>; Yb<sup>+</sup><sup>3</sup>; Lu<sup>+</sup><sup>3</sup>; Ti<sup>+</sup><sup>4</sup>; Zr<sup>+</sup><sup>4</sup>; Ti<sup>+</sup><sup>3</sup>; Mf<sup>+</sup><sup>4</sup>; Nb<sup>+</sup><sup>5</sup>; Ta<sup>+</sup><sup>5</sup>; Nb<sup>+</sup><sup>4</sup>; Ta<sup>+</sup><sup>4</sup>, V<sup>+</sup><sup>5</sup>; V<sup>+</sup><sup>4</sup>; Nb<sup>+</sup><sup>3</sup>; Mn<sup>+</sup><sup>3</sup>; Mn<sup>+</sup><sup>4</sup>; Mn<sup>+</sup><sup>3</sup>; Mn<sup>+</sup><sup>4</sup>; Pe<sup>+</sup><sup>2</sup>; Fe<sup>+</sup><sup>3</sup>; Co<sup>+</sup><sup>2</sup>; Co<sup>+</sup><sup>3</sup>; Ni<sup>+</sup><sup>2</sup>; Ni<sup>+</sup><sup>3</sup>; Ni<sup>+</sup><sup>4</sup>; Ru<sup>+</sup><sup>2</sup>; Ru<sup>+</sup><sup>3</sup>; Ru<sup>+</sup><sup>4</sup>; Rh<sup>+</sup><sup>3</sup>; Ir<sup>+</sup><sup>3</sup>; Rh<sup>+</sup><sup>2</sup>; Ir<sup>+</sup><sup>2</sup>; Pd<sup>+</sup><sup>4</sup>, Pt<sup>+</sup><sup>4</sup>; Pd<sup>+</sup><sup>2</sup>; Pt<sup>+</sup><sup>2</sup>; Os<sup>+</sup><sup>4</sup>; Cu<sup>+</sup>; Cu<sup>+</sup>; Cu<sup>+</sup><sup>2</sup>; Cu<sup>+</sup><sup>3</sup>; Ag<sup>+</sup>; Ag<sup>+</sup><sup>2</sup>; Ag<sup>+</sup><sup>3</sup>; Au<sup>+</sup>; Au<sup>+</sup><sup>3</sup>; Au<sup>+</sup><sup>3</sup>; Au<sup>+</sup><sup>3</sup>; Zh<sup>+</sup><sup>2</sup>; Cd<sup>+</sup><sup>2</sup>; Hg<sup>+</sup>; Hg<sup>+</sup><sup>2</sup>; Al<sup>+</sup><sup>3</sup>; Ga<sup>+</sup>; In<sup>+</sup><sup>3</sup>; In<sup>+</sup>; Tl<sup>+</sup><sup>3</sup>; Tl<sup>+</sup>; Ge<sup>+</sup><sup>4</sup>; Ge<sup>+</sup><sup>2</sup>; Sn<sup>+</sup><sup>4</sup>; Sn<sup>+</sup><sup>2</sup>; Pb<sup>+</sup><sup>4</sup>; Pb<sup>+</sup><sup>2</sup>; Sb<sup>+</sup><sup>3</sup>; Sb<sup>+</sup><sup>5</sup>; As<sup>+</sup><sup>3</sup>; As<sup>+</sup><sup>3</sup>; Bi<sup>+</sup><sup>5</sup>; organic compounds containing at least one N<sup>+</sup> site; organic compounds containing at least one phosphonium site; organic compounds containing at least one selenonium site;

178. (Previously Presented) The conversion coating of claim 177 wherein the cationic solubility control agent is selected from  $H^{\dagger}$ ;  $Li^{\dagger}$ ;  $Na^{\dagger}$ ;  $K^{\dagger}$ ;  $Rb^{\dagger}$ ;  $Cs^{\dagger}$ ;  $NH_4^{\dagger}$ ;  $Mg^{\pm 2}$ ;  $Ca^{\pm 2}$ ;  $Sr^{\mp 2}$ ;  $Y^{\pm 3}$ ;  $La^{\pm 3}$ ;  $Ce^{\pm 3}$ ;  $Ce^{\pm 4}$ ;  $Nd^{\pm 2}$ ;  $P^{\pm 3}$ ;  $P^{$ 

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stibonium site; organic compounds containing at least one oxonium site; organic compounds containing at least one sulfonium site; organic compounds containing at least one iodonium site; quaternary ammonium compounds having a formula NR<sub>4</sub><sup>+</sup>, where R is an alkyl, aromatic, or acvelic organic constituent; or combinations thereof.

179. (Previously Presented) The conversion coating of claim 165 wherein the conversion coating is colored.

180. (Previously Presented) The conversion coating of claim 179 further comprising an agent which improves color-fastness of the conversion coating.

181. (Previously Presented) The conversion coating of claim 180 wherein the agent which improves color-fastness is selected from an active UV blocker, a passive UV blocker, a brightener, or a combination thereof.

182. (Previously Presented) The conversion coating of claim 181 wherein the agent which improves color-fastness is the active UV blocker selected from carbon black, graphite, phthalocyanines, or combinations thereof.